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***In-Vitro* Antimicrobial Susceptibility and Phytochemical Constituents of Methanol Leaf Extract of *Prosopis africana* against some Selected Microorganisms.**

Abstract

The idea that certain plants had healing potential was known long before human beings discovered the existence of pathogens. The crude methanolic leaf extract of *Prosopis africana* was assayed for antimicrobial potency using Agar-well diffusion technique against *Salmonella typhi*, *Klebsiella pneumoniae*, *Streptococcus pyogenes*, *Pseudomonas aeruginosa*, Methicillin-Resistant *Staphylococcus aureus* (MRSA) and *Candida albicans*. Qualitative phytochemical screening was also carried out. The result of the antimicrobial screening showed antimicrobial potency against the test isolates with various degrees of zone of inhibition which varied between 10mm – 22mm. The highest zone was noted against *Klebsiella pneumoniae* (22mm), followed by *Streptococcus pyogenes* and *Candida albicans* (21mm), *Salmonella typhi* (20mm), MRSA (19 mm) then *Pseudomonas aeruginosa* (18mm). Chloramphenicol and Fluconazole are used as reference standard and their zones of inhibitions ranged from 26mm–29mm. The Minimum Inhibitory Concentration (MIC) of the extract ranged between 12.50mg/mL – 50.00mg/mL whilst the Minimum Bactericidal Concentration (MBC) and Minimum Fungicidal Concentration (MFC) of the extract were at 50.00mg/mL. The result of phytochemical screening revealed the presence of carbohydrates, flavonoids, phenols/tannins, saponins, terpenes, steroids, cardiac glycosides and alkaloids as secondary metabolites. The result of this study showed that methanolic leaf extract of *P. africana* has proved its use in folklore as an alternative

24 antimicrobial agent and further research can lead to isolation of a new lead of medical
25 importance.

26 **Key words:** Antimicrobial Susceptibility, *Prosopis africana*, Phytochemical Constituents, 96-
27 well microplates, MIC, MBC and MFC.

28

29 **1. Introduction**

30 Infectious diseases are particularly a major challenge to public health, despite tremendous
31 scientific discovery of medicines for their treatment [1]. This is due to increase in resistance to
32 antibiotics by pathogenic microorganisms as a result of misuse and over prescription of
33 antibiotics which has affected our ability to treat patients empirically [2][3]. So in recent years
34 there has been continuous and urgent need to discover new antimicrobial compounds with
35 diverse chemical structures and novel mechanisms of action because the incidence of new and
36 re-emerging infectious diseases and development of resistance to the antibiotics in current
37 clinical use [4]. However, nature is endowed with providing continuous new biomolecules with
38 novel structures that are designed to interact with biological systems to provide defense against
39 infectious diseases [5].

40 The presence of bioactive principles such as alkaloids, phenols, tannins, glycosides and essential
41 oils amongst others is responsible for the medicinal properties of plants [6]. It is necessary to
42 screen medicinal plants for the presence of these bioactive chemicals which may lead to a new
43 active principle. Scientific studies available on medicinal plants indicate that promising
44 phytochemical can be developed for many health problems [7]. The benefits of using plant
45 derived medicines are that they are relatively cheaper and stable.

46 *P. africana* (Guill. & Perr.) Taub. belongs to kingdom plantae, division tracheophyta, class
47 magnoliopsida, order fabales and family fabaceae. It is a renowned and versatile tree of immense
48 economic value amid the rural communities in the Guinea savanna of Nigeria. It is mostly found
49 in savanna countries of Africa like Senegal and Nigeria [8]. Literature gives an account of its
50 uses as folk medicines for several ailment and virtually all its parts are of medicinal value. [9]
51 reported that the stem bark is used as remedies for dysentery, gonorrhoea, bronchitis and skin
52 diseases. In Niger State of Nigeria, the twigs, leaves, bark, and secondary roots are used for
53 treatment and relieve of typhoid fever, dental decay, malaria as well as stomach cramps while,
54 [10] attested that the bark and root decoctions are utilized for the treatment of Trypanosomiasis
55 in cattle and on lesions as a lotion.

56 It is in knowledge domain that antimicrobial resistance is a great challenge to holistic treatment
57 of infectious diseases as a result of either the use of substandard antibiotics, misuse or over
58 prescription. There is therefore, need for search to discovering new lead principles that will be
59 effective, safe, readily available and cost effective. This study was aimed to determine the
60 antimicrobial properties of *P. africana* towards the development of new antimicrobial agent.

61 **2.0 Materials and Methods**

62 **2.1 Plant Sample Collection and Identification**

63 The leaves of *P. africana* was collected from Bida in Niger State, Nigeria, voucher specimen
64 was prepared, the plant was identified and voucher specimen was deposited in herbarium unit of
65 the Department of Medicinal Plant Research and Traditional Medicine (MPR&TM), National
66 Institute for Pharmaceutical Research and Development (NIPRD), Idu-Abuja, Nigeria.

67 **2.2 Preparation and Preservation of Plant Material**

68 The leaves were properly washed under clean-running tap water to remove the dirt and air dried
69 at room temperature for a week. The dried leaves were pulverized into powder with clean
70 wooden pestle and mortar, and sieved. The pulverized sample was stored in a clean plastic
71 container, properly labeled and tightly covered at 37°C prior for further analysis.

72 **2.3 Extraction of Crude Extract**

73 One hundred grams (100g) of the pulverized leave of *P. africana* was accurately weighed and
74 subjected to cold maceration in 500mL of absolute methanol for 72 h at laboratory temperature.
75 The macerated extract was filtered using Whatman No.1 filter paper. The extraction was
76 repeated for the maximum extraction of the active ingredients and to also obtain reasonable
77 yield (crude extract). The filtrate was dried using water bath at 45°C until all the solvent
78 evaporated out.

79 The percentage yield of the crude extract (PYCE) was calculated using the formula by [11].

$$80 \quad PYCE = \frac{\text{Mass of the crude extract obtained}}{\text{Mass of the pulverized plant sample}} \times 100$$

81

82 **2.4 Phytochemical Screening of the Extract**

83 The qualitative phytochemical screening of the leaves of *P. africana* was carried out in
84 Pharmacognosy unit, Department of Medicinal Plant Research and Traditional Medicine,
85 NIPRID, Idu-Abuja. The standard methods illustrated by [12][13] [14] were adopted to test for
86 the presence of carbohydrates, flavonoids, phenols/tannins, saponins, terpenes, steroids, alkaloids
87 and cardiac glycosides.

88 **3.4.1 Determination of carbohydrates: (Molisch's Reagent Test)**

89 The fraction of the extract was mixed with 2mL of Molisch's reagent and the mixture was
90 properly shaken. Thereafter, 2mL of concentrated H₂SO₄ was slowly and carefully added. The
91 appearance of a purple ring/violet at the interphase signifies the presence of carbohydrates.

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93 **3.4.2 Determination of flavonoids**

94 **i. Alkaline Reagent Test**

95 To the extract, 2mL of 2% solution of NaOH was mixed. An intense yellow colouration was
96 formed which later turned colourless when few drops of diluted acid was added which signifies
97 the presence of flavonoids.

98 **ii. Shinoda Test**

99 To the extract, few drops of Mg ribbon fragments were added followed by concentrated HCl.
100 Pink scarlet colour appeared after few minutes which signifies the presence flavonoids.

101 **3.4.3 Determination of phenols/tannins**

102 The fraction of the extract was mixed with 2mL of 2% solution of FeCl₃. A blue-green or black
103 colouration which appeared signifies the presence of phenols and tannins.

104 **3.4.4 Determination of saponins: (Froth Test)**

105 The fraction of the extract was mixed with 5mL of distilled H₂O in a test tube and was
106 vigorously shaken for 30 seconds. The formation of the stable foam signifies the presence of
107 saponins.

108 **3.4.5 Determination of terpenes: (Liebermann's Test)**

109 The fraction of the extract was mixed with each of 2mL of chloroform and 2mL of acetic acid.
110 The mixture was then cooled down. Thereafter, concentrated H₂SO₄ was slowly and carefully
111 added and the colour changed from violet to blue which signifies the presence of terpenes.

112 **3.4.6 Determination of steroids: (Salkoski's Test)**

113 To the extract, about 2mL of the chloroform was mixed. Concentrated H₂SO₄ was slowly and
114 carefully added and shaken gently. A reddish brown colour signifies the presence of steroids.

115 116 **3.4.7 Determination of cardiac glycosides: (Keller-kilani Test)**

117 The extract was mixed with 2mL of glacial acetic acid containing 1-2 drops of 2% solution of
118 FeCl₃. The mixture was then poured into another test tube containing 2mL of concentrated
119 H₂SO₄. A brown ring at the interphase signifies the presence of cardiac glycosides.

120 **3.4.8 Determination of alkaloids**

121 **i. Dragendoff's Test**

122 To 2mL of 1% HCl, the extract was added. Dragendoff's reagent was then added to the mixture.
123 The resulting turbidity of the precipitate was considered as a proof for the presence of alkaloids.

124 **ii. Hagner's Test**

125 To another 2mL of 1% HCl, the extract was added. Hagner's reagent was then added to the
126 mixture. The resulting turbidity of the precipitate was deemed as an evidence for the attendance
127 of alkaloids.

128 **iii. Wagner's Test**

129 To another 2mL of 1% HCl again, the extract was added. Wagner's reagent was then added to
130 the mixture. The resulting turbidity of the precipitate was believed to be a testimony for the
131 existence of alkaloids.

132 **2.5 The Test Microorganisms**

133 Antimicrobial activity of methanol extract of leaves of *P. africana* was investigated against five
134 bacterial isolates and one fungal isolate which were obtained from Vaccine Discovery and
135 Research Laboratory, Centre for Genetic Engineering and Biotechnology, Federal University of
136 Technology, Minna, Nigeria. The bacteria strains which were for the study include *Salmonella*
137 *typhi*, *Klebsiella pneumoniae*, *Streptococcus pyogenes*, *Pseudomonas aeruginosa*, Methicillin-
138 Resistant *Staphylococcus aureus* (MRSA) and the fungi used for the study was *Candida*
139 *albicans*. The tested bacteria were maintained on Nutrient agar at 37°C for 24 h and *Candida*
140 *albicans* on Potatoes Dextrose Agar at 30°C for 48-72 h.

141 **2.6 Inoculum Preparation**

142 A loopful of isolated colonies were inoculated into 4 mL sterile Mueller-Hinton Broth (MHB)
143 for bacteria and Sabouraud Dextrose Broth (SDB) for fungi and incubated at 37°C for 2 h. The
144 turbidity of actively growing microbial suspensions were adjusted with freshly prepared MHB
145 and SDB using BaSO₄ turbidity standard to match turbidity standard of 0.5 McFarland. This
146 turbidity was equivalent to approximately 1.5x10⁸ CFU/mL cells for bacteria, and 1.5x10⁷
147 spores/mL for fungi strain. The grown suspension was used for further testing.

148 **2.7 Preparation of Crude Extract**

149 For the preparation of the stock solution, 0.5g of the crude extract was accurately weighed using
150 analytical weighing balance into a sterile tube containing 1mL of 2% Dimethylsulfoxide
151 (DMSO). This was vortexed to allow the extract to completely dissolve and 9mL of sterile
152 distilled water was added to give final extract concentration of 50mg/mL. Double fold dilutions
153 was carried out to give extract concentrations of 25mg/mL, 12.5mg/mL and 06.25mg/mL using
154 sterile distilled water respectively.

155 **2.8 *In-vitro* Antimicrobial Susceptibility Assay of the Extract**

156 Susceptibility test of the extract against the isolates were determined in the Microbiology
157 Laboratory, Ibrahim Badamasi Babangida University Lapai in Niger State, Nigeria using Kirby-
158 Bauer agar diffusion method according to NCCLS standards ([15][16]. The Mueller-Hinton Agar
159 (MHA) and Sabouraud Dextrose Agar (SDA) were used for the antimicrobial activity test. About
160 100 μL of MHB and SDB cultures containing 0.5 McFarland equivalent to approximately
161 1.5×10^8 CFU/mL cells for bacteria, and 1.5×10^7 spores/mL for fungi strain were dispensed into
162 empty sterile petri dishes using micropipettes. Twenty three millilitres (23mL) of sterilized MHA
163 and SDA maintained between $50 - 45^\circ\text{C}$ was added to the appropriate petri dishes and rocked
164 gently for even distribution of the organisms under aseptic condition and allowed to gel under
165 safety hood for 1 h. On each of the plates containing bacteria isolates. Five wells of 8 mm in
166 diameter were made on the agar plates using sterile metallic cork borer and labelled properly.
167 The base of the wells was sealed with 30 μL of MHA and SDA. Thereafter, 200 μL of different
168 concentrations of the extract were carefully and aseptically added with the aid of micropipette
169 into each well and left in the safety hood for 2 h for proper diffusion of the extracts into the agar
170 and then incubated at 37°C for 24 h for bacteria. The same procedure was repeated for fungi
171 strain and incubated at 25°C for 48 h for fungi. The experiment was set up in duplicates. The
172 plates were observed for activity and zones of inhibitions were measured and recorded as mean
173 zone of inhibition. The diameter of each zone was accurately measured with a spotless and
174 translucent ruler in millimetre (mm) in vertical and horizontal manner and mean was determined.

175 Control experiments were set up by using standard antibiotics, Chloramphenicol (250mg) for
176 bacteria strain and fluconazole (80mg) for fungi specie as reference standards for positive
177 control. Sterile MHA and SDA plates were used as Media Sterility Control (MSC) and MHA and

178 SDA plates with the used organisms streaked as Organism Viability Control (OVC). All the
179 controls were given the same treatment as the experiments.

180 **2.9 Determination of Minimum Inhibitory Concentration (MIC)**

181 The minimum inhibitory concentration (MIC) value of the extract of *P. africana* leaves extract
182 was determined by microdilution broth method in 96-well microplates [17]. Chloramphenicol
183 (Fidson, Nigeria) and Fluconazole (Pfizer, UK) were used as the standard drug for bacteria and
184 fungi at stock concentration of 50µg/ml. Controls of sterility for the Mueller-Hinton nutrient
185 broth, control culture (inoculum), Chloramphenicol, Fluconazole, crude extract and DMSO were
186 carried out. The microwell plates were closed and incubated aerobically at 37°C for 24 h.
187 Thereafter, 50µL of tetrazolium dye was applied into each well with further 2 h incubation at
188 37°C and colour change was observed. Any well with reddish-pink colour signifies the microbial
189 growth, which was noted and documented as positive (MIC). All assays were carried out in
190 triplicate.

191 MIC was carried out on the extract against the isolates using the broth microdilution method
192 (BMM) in 96 micro wells titre plates using NCCLS method [16] with little modification. A
193 volume of 50µL of the extract was dispensed into first row and the same volume of the sterilized
194 media (MHB and SDB) was dispensed into each well except the first row. A two-fold dilution
195 was carried out from row 2 by taking 50µL of the extract to the next row, mixed well and the
196 serial dilution continued to row 7 where 50µL from the wells was discarded away. Then, 50µL
197 of 0.5 McFarland of 2 h culture was added to each well in row 1-7. The rows 8 and 9 were the
198 OVC and MSC. The plates were incubated at 37°C for 24 h. The test was carried out in duplicate
199 and the values are express in mean.

200 **2.10 Determination of Minimum Bactericidal and Fungicidal Concentration (MBC and** 201 **MFC)**

202 The MBC and MFC were determined from the Minimum Inhibitory Concentration (MIC) result
203 by subculturing from the wells that shows no any sign of turbidity in the MIC test and streak on
204 the freshly prepared MHA and SDA plates and incubate at 37°C for 24 to 48 h and the plates
205 were checked for the present or absent of the growth.

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208 **3.0 Results**

209 **3.1 Phytochemical Constituents**

210 Qualitative biological active compounds of the methanolic crude extract disclosed the occurrence
211 of carbohydrates, flavonoids, phenols/tannins, saponins, terpenes, steroids, cardiac glycosides
212 and alkaloids respectively as demonstrated in table 1 below.

213 **3.2 Antimicrobial Activity**

214 The methanolic crude extract has antimicrobial efficacy against all the test isolates at 50 and 25
215 mg/mL of concentration, while *Salmonella typhi*, *Klebsiella pneumoniae*, *Streptococcus*
216 *pyogenes* and Methicillin-Resistant *Staphylococcus aureus* (MRSA) showed activity even at
217 12.50mg/mL. The extract at 0.625mg/mL however had no activity against all the tested
218 organisms. The antimicrobial controls were active against all tested organisms except
219 *Pseudomonas aeruginosa* (figure 1).

220 **3.3 Minimum Inhibitory and Bactericidal Concentrations of the Crude Extract against** 221 **Sensitive Organisms**

222 Minimum Inhibitory Concentration (MIC) was carried out on all the isolates where *Salmonella*
223 *typhi* and *Streptococcus pyogenes* had MIC of 12.50mg/mL. MRSA, *Klebsiella pneumoniae* and

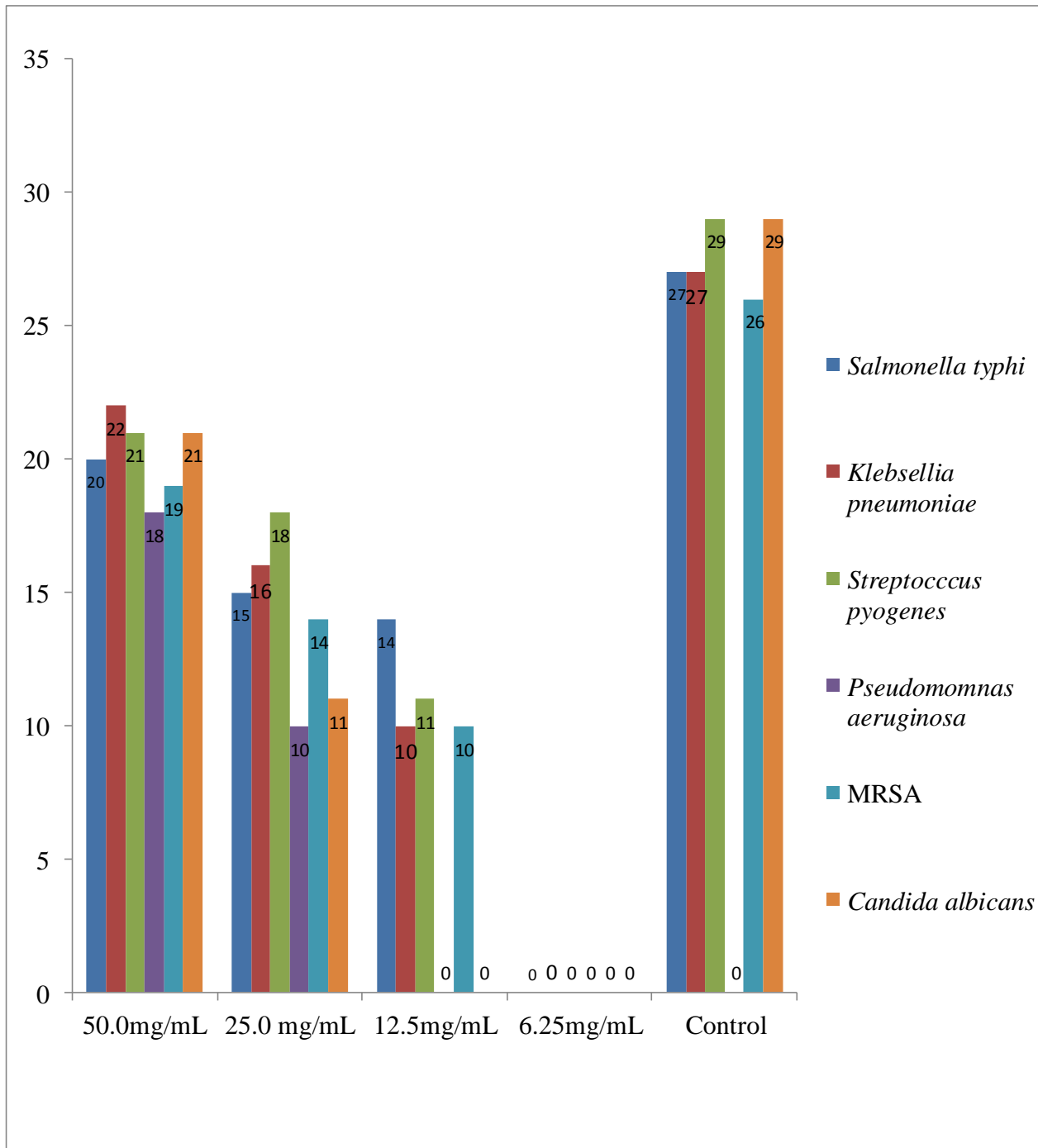
224 *C. albicans* had MIC of 25mg/mL while the MIC of *Pseudomonas aeruginosa* was at 50mg/mL
 225 (table 2). The MBC of *Salmonella typhi*, *Klebsiella pneumoniae*, *Streptococcus pyogenes*,
 226 MRSA were at 50mg/mL and *C. albicans* had the MFC of 50mg/mL (table 3).

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Table1: Phytochemical constituents of methanolic leaves

Phytochemicals	Tests	Inference
Carbohydrates	Molisch	+
Flavonoids	Alkaline	+
Phenol/Tannins	Ferric Chloride	+
Saponins	Froth	+
Terpenes	Liebermann	+
Steroids	Salkowski	+
Alkaloids	Dragendoff's	+
	Hagner's	+
	Wagner's	+
Cardiac glycosides	Keller-Kilani	+

233 Key: + = Present - = Absent



234 Fig. 1: Mean zone of inhibition of methanolic extract of *P. africana* on the test organisms (mm).

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238 **Table 2: Minimum Inhibitory Concentration of Methanol Extract of *P. africana* on the test**
 239 **Organisms**

Organisms	Minimum Inhibitory Concentration (mg/mL)	Control (mg/mL)
<i>Salmonella typhi</i>	12.5	50
<i>Streptococcus pyogenes</i>	12.5	50
MRSA	25.0	50
<i>Klebsiella pneumoniae</i>	25.0	50
<i>Pseudomonas aeruginosa</i>	50.0	50
<i>Candida albicans</i>	25.0	50

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241 **Table 3: Minimum Bactericidal and Fungicidal Concentration of Methanol Extract of *P. africana***
 242 **on the test Organisms**

Organisms	MBC/MFC (mg/mL)
<i>Salmonella typhi</i>	50.0
<i>Streptococcus pyogenes</i>	50.0
MRSA	50.0
<i>Klebsiella pneumoniae</i>	50.0
<i>Candida albicans</i>	50.0

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244 **4.0 Discussion**

245 Determining the phytochemical constituents to identify the possible therapeutic agents contained
 246 in this plant in order to establish the basis for their uses in folklore medical practice is important
 247 to discovery of new active principle.

248 Variety of different natural chemical compounds such as saponins, tannins, alkaloids, terpenoids,
249 cyanoglycosides, oleic, flavonoids and stearic acids have been found to confer antimicrobial
250 properties to plants and plant products [18] [19] [20]. [21] stated that flavonoids are polyphenolic
251 phytocompounds which are found in plants and possess antifungal, antibacterial, anticancer, anti-
252 inflammatory as well as antioxidant properties. In accordance with the study conducted by [22]
253 the seed and pod of *Prosopis africana* shows the presence of saponin, alkaloids, steroids,
254 flavonoids, phlabotanin and tannin, the result of this study also showed the presence of saponin,
255 alkaloids, steroids, flavonoids, carbohydrates, terpenes, cardiac glycosides and tannin which is in
256 agreement with findings by [6] but different from the findings by [23], where tannins, saponins
257 and terpenes were absent in methanol leaf extract of *Prosopis africana*. The study by [24],
258 showed the absence of tannins in *Prosopis lappacea*. The occurrence of these bioactive
259 compounds in reasonable amount in the leaves of *P. africana* could have been responsible for its
260 characteristic antimicrobial properties. These are recognized to have antibacterial agent and may
261 be utilized traditionally for the treatment of infectious diseases [25]. [21] stated that flavonoids
262 are polyphenolic phytocompounds which are found in plants and possess antifungal,
263 antibacterial, anticancer, anti-inflammatory as well as antioxidant properties.

264 Antimicrobial analysis of methanol crude leaf extract of *P. africana* exhibited some level of
265 antimicrobial properties against microorganisms which include *Streptococcus pyogenes*,
266 *Pseudomonas aeruginosa*, Methicillin-Resistant *Staphylococcus aureus* (MRSA) *Salmonella*
267 *typhi*, *Klebsiella pneumoniae*, and the fungi used for the study was *Candida albicans* at various
268 concentrations from 50mg/mL to 6.25mg/mL with varied diameters zones of inhibition ranging
269 from 1mm for 29mm. [22] tested the aqueous and methanol extract of seed and pod of *P.*
270 *africana* against fifteen bacteria at a fix concentration of 25mg/mL and recorded zone of

271 inhibition ranging from 5mm to 17mm. [26] also reported the antimicrobial activities of *P.*
272 *africana* ethanol and aqueous stem and root extract against clinical isolates of oral pathogens
273 namely *Streptococcus mutans*, *Staphylococcus saprophyticus* and *C. albicans* with significant
274 action. The study by [25], revealed activity against *Pseudomonas aeruginosa*, *Staphylococcus*
275 *aureus*, and *Bacillus subtilis* at concentrations comparable to the findings of this study. However,
276 the study by [27], showed no activity against *K. pneumoniae*. The result of this study showed no
277 activity against tested organisms at concentration of 6.25mg/mL. Interestingly, the extract had
278 activity against MRSA at 12.5mg/mL concentration with MIC of 25.0mg/mL. This is very
279 promising because of the facts that further purification may exhibit better activity which could
280 lead to discovery of a new lead against antibiotics resistant *Staphylococcus aureus*. It is also
281 worthy of notice that the extract is also active against *Pseudomonas aeruginosa* at crude MIC of
282 50.0mg/mL.

283 **5. Conclusion**

284 The methanolic leaf extract of *P. africana* has displayed varied activity against pathogenic
285 microorganisms and could represent candidate of antimicrobial agent against some human
286 pathogenic microbes. Furthermore, the bioactive ingredients indicated that the plant part have
287 proved its usage in the folkloric medicine for the management of different ailments and could be
288 the basis of alternative anti-infective therapy. Therefore, these findings shall broaden and
289 enhance global data base of the antimicrobial property of the active ingredients present.

290 **6. Limitation of the study**

291 The limitation of this study includes lack of funding to permit us to buy solvents for
292 fractionations which will enable us to obtain a pure compound.

293 **Conflict of Interest**

294 The authors declared no conflict of interest.

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